(US)
Access DB# 90/47

SEARCH REQUEST FORM

7.502 (77)

Scientific and Technical Information Center

Requester's full Name:_	Devesh Khare Examiner #:_	77931	Date:	3/27/2003	
Art Unit: 1623	Phone Number <u>605-1199</u>			9/775,760	•
Mail Box: <u>CM1-8B19</u> and BI MAIL	dg/Room Location: <u>CM1-8A13</u> Re	esults Format	Preferred	(circle):PAPER	DISK E-
If more than one searcl	1 is submitted, please prioriti ***********************************	ze searche ******	s in ord	er of need.	****
Please provide a detailed statem search include the elected spec- the concept or utility of the inve	nent of the search topic, and describe a ies or structures, key words, synonyms ention. Define any terms that may hav . Please attach a copy of the cover she	s specifically , acronyms, a e a special me	as possible nd registry caning. Gi	e the subject mat numbers, and co	ter to be
Title of Invention: See B	ib Data Sheet		·		. 1
Inventors (please provide ful	I names): See Bib Data Sheet				
				, · · · ·	
Earliest priority Filing Da	ate: See Bib Data Sheet				
	Please include all pertinent information	on (parent, ch	ild, divisio	onal, or issued pa	tent
Please carry out a	search for a process for produc	cing polysa	ccharide	es ethers (clair	me 33_
	lucing cellulose ethers (claims				
The Bib Data Shee	et which discloses the inventor	names, tit	le of the	invention, an	d the
earliest priority filing date					
			Tec	Mary Jane Re h. Info. Speciall TC-1600	ihi st, STIC
Thank you.	py of the claims with the search	h.	(CM-1, Room 6/ Phone: 605-1	\-06 155
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STAFF USE ONLY	Type of Search	Vendors	and cost	where applicable	è
Searcher:		STN		<u>-</u>	
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Smenter Search

Khare 09/775,760

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L25 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

2003:43049 HCAPLUS

DOCUMENT NUMBER:

138:91690

TITLE:

Recycling of salts in the manufacture of modified

polysaccharide ethers from sodium salt stock

INVENTOR(S):

Mallon, Charles B.; Vames, John S. ; Sarlis, John I.; See, Benito; Trampe, David M.; Datta, Rathin

USA

PATENT ASSIGNEE(S): SOURCE:

U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 2003013871 20030116 US 2001-775760 20010202

PRIORITY APPLN. INFO.: US 2001-775760 20010202 Salts formed during the manuf. of polysaccharide ethers, e.g., sodium nitrate or sodium acetate, are converted to their corresponding acids and bases by means of an elec. current, preferably in combination with a bipolar membrane and suitable cation and/or anion membranes. The acids and bases recovered from the processes can be recycled, thereby avoiding the need to provide for disposal of the salts. Thus, a stream contg. .apprx.8% Na acetate, 0.5% NaOH, 4% glycols, and 0.5% cellulosic material recovered from a synthesis of hydroxyethyl cellulose was processed in a TS-2 ED stack contg. 6 cell pairs made up of AM-1, CM-2 and CURIUM ion-exchange membranes. The pH of the feed was raised to .apprx.11.5 by addn. of 40% NaOH whenever the pH dropped to .apprx.11. As a result of this pH control, .apprx.91% of the acetate was transferred and cell resistance was kept under control. Na acetate concn. in the conc. was .apprx.20%. The product from the ED step was then processed through a column contg. Duolite C-467 to remove multivalent cations. Ca and Mg cations were below 1 ppm in the resulting stream. This product stream was then subjected to water splitting electrodialysis using a 2-compartment TS-2 stack equipped with BP1, CM-1, and CMX ion exchange membranes. The stream processed with no difficulty and produced an acid/salt product contg. about 16% acetic acid and a sodium hydroxide product with a concn. of .apprx.10%. There was no evidence of irreversible membrane fouling and cell voltage remained low (.apprx.1.2 V/cell pair) throughout the run.

ICM C08B011-00

TCS C08B011-12

536084000

43-3 (Cellulose, Lignin, Paper, and Other Wood Products) Section cross-reference(s): 44

ST cellulose acetate salt recycling electrolysis bipolar membrane

IT Membranes, nonbiological

> (bipolar; method for recycling of salts in manuf. of modified cellulose ether)

IT Polysaccharides, preparation

RL: IMF (Industrial manufacture); PREP (Preparation)

(ethers; method for recycling of salts in manuf. of modified cellulose ether)

IT Electrodialysis

```
Ion exchange membranes
        (method for recycling of salts in manuf. of modified cellulose ether)
IT
     Acids, preparation
     Bases, preparation
     RL: IMF (Industrial manufacture); PUR (Purification or recovery); PREP
     (Preparation)
        (regeneration of; method for recycling of salts in manuf. of modified
        cellulose ether)
     1398-61-4DP, Chitin, derivs.
IT
                                    9000-01-5DP, Gum Arabic, derivs.
     9000-07-1DP, Carrageenan, derivs.
                                        9000-30-0DP, Guar, derivs.
     9000-65-1DP, Tragacanth, derivs.
                                        9000-69-5DP, Pectin, derivs.
     9002-18-ODP, Agar, derivs.
                                  9004-24-4P, Carboxymethyl cellulose aluminum
            9004-34-6DP, Cellulose, derivs.
                                              9004-62-0P, Hydroxyethyl
                 9005-25-8DP, Starch, derivs.
     cellulose
                                                9005-32-7DP, Alginic acid,
              9012-76-4DP, Chitosan, derivs.
                                                9050-04-8P, Carboxymethyl
     cellulose calcium salt
                              9086-60-6P, Carboxymethyl cellulose ammonium salt
     11138-66-2DP, Xanthan gum, derivs.
                                          55962-76-0P, Carboxymethyl cellulose
     lithium salt
                    66988-45-2P, 2-Sulfoethyl cellulose magnesium salt
                    124363-51-5P, Carboxymethyl sulfoethyl cellulose calcium
     124363-50-4P
     salt
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (method for recycling of salts in manuf. of modified cellulose ether)
IT
     1310-65-2P, Lithium hydroxide
                                    9000-11-7P, Carboxymethyl cellulose
     RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT
     (Reactant or reagent)
        (method for recycling of salts in manuf. of modified cellulose ether)
IT
     9088-04-4, Carboxymethyl hydroxyethyl cellulose sodium salt
     RL: PEP (Physical, engineering or chemical process); PYP (Physical
     process); RCT (Reactant); PROC (Process); RACT (Reactant or reagent)
        (method for recycling of salts in manuf. of modified cellulose ether)
     62-54-4, Calcium acetate
                               139-12-8, Aluminum acetate
                                                            142-72-3,
     Magnesium acetate
                        7664-41-7, Ammonia, reactions 9004-32-4,
     Carboxymethyl cellulose sodium salt 9032-46-6, Sulfoethyl cellulose
     39277-57-1, Sulfoethyl cellulose sodium salt 117989-25-0, Carboxymethyl
     sulfoethyl cellulose sodium salt
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (method for recycling of salts in manuf. of modified cellulose ether)
     75-21-8, Ethylene oxide, reactions
                                         75-56-9, Propylene oxide, reactions
     26249-20-7, Butylene oxide
     RL: RGT (Reagent); RACT (Reactant or reagent)
        (method for recycling of salts in manuf. of modified cellulose ether)
IT
     59249-79-5, Aciplex K 101
     RL: TEM (Technical or engineered material use); USES (Uses)
        (method for recycling of salts in manuf. of modified cellulose ether)
     64-19-7P, Acetic acid, preparation 1305-62-0P, Calcium hydroxide,
    preparation
                 1309-42-8P, Magnesium hydroxide 1310-58-3P, Potassium
     hydroxide, preparation 1310-73-2P, Sodium hydroxide, preparation
     1336-21-6P, Ammonium hydroxide 7647-01-0P, Hydrochloric acid,
    preparation
                  7664-38-2P, Phosphoric acid, preparation 7664-93-9P,
     Sulfuric acid, preparation 7697-37-2P, Nitric acid, preparation
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (regeneration of; method for recycling of salts in manuf. of modified
        cellulose ether)
L25 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                         2000:117084 HCAPLUS
DOCUMENT NUMBER:
                         132:153513
TITLE:
                         Production of polysaccharide hydroxyalkyl ethers
```

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INVENTOR(S):
                          Mallon, Charles B.; Vames, John S.
                          ; Sarlis, John Ioannis; See, Benito
                          ; Trampe, David M.; Datta, Rathin
PATENT ASSIGNEE(S):
                          Union Carbide Chemicals & Plastics Technology Corp.,
SOURCE:
                          PCT Int. Appl., 32 pp.
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                       KIND DATE
                                            APPLICATION NO. DATE
     WO 2000008059
                             20000217
                       Al
                                           WO 1999-US17597 19990803
         W: AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GE, HU, ID, IL, IS,
              JP, KR, KZ, LC, LK, LR, LT, LV, MG, MK, MN, MX, NO, NZ, PL, RO,
             RU, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY,
            . KG, KZ, MD, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                           20000217
     CA 2339374
                       AA
                                           CA 1999-2339374 19990803
     AU 9954643
                       A1
                             20000228
                                           AU 1999-54643
                                                             19990803
     EP 1109834
                       A1
                             20010627
                                           EP 1999-940869
                                                             19990803
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
     BR 9912637
                       Α
                            20011009
                                            BR 1999-12637
                                                             19990803
PRIORITY APPLN. INFO.:
                                         US 1998-95263P
                                                         P 19980804
                                         WO 1999-US17597 W 19990803
     Processes for the prodn. of polysaccharide ethers, e.g. cellulose
     hydroxyethyl ether, are disclosed wherein salts formed after the swelling
     and neutralization of the process, e.g., sodium nitrate or sodium acetate,
     are converted to their corresponding acids and bases by means of an elec.
     current, preferably in combination with a bipolar membrane and suitable
     cation and/or anion membranes. The acids and bases recovered from the
     processes can be recycled, thereby avoiding the need to provide for
     disposal of the salts.
IC
     ICM C08B011-20
CC
     43-3 (Cellulose, Lignin, Paper, and Other Wood Products)
     cellulose ether manuf salt recycling; bipolar membrane salt deionization
     acid recycling; electrodialysis salt sepn cellulose etherification
     Membranes, nonbiological
IT
        (bipolar; electrodialysis of salts after prodn. of polysaccharide
        hydroxyalkyl ethers)
IT
     Electrodialysis
     Etherification
     Ion exchange membranes
        (electrodialysis of salts after prodn. of polysaccharide hydroxyalkyl
        ethers)
IT
     Polysaccharides, preparation .
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (ethers; electrodialysis of salts after prodn. of polysaccharide
        hydroxyalkyl ethers)
IT
     9004-62-0P, Cellulose hydroxyethyl ether
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (electrodialysis of salts after prodn. of polysaccharide hydroxyalkyl
        ethers)
```

IT 125935-42-4, Duolite C 467

RL: TEM (Technical or engineered material use); USES (Uses) (ion exchanger; electrodialysis of salts after prodn. of polysaccharide hydroxyalkyl ethers)

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> log hold .		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
FULL ESTIMATED COST	ENTRY	SESSION
TOUR EDITINIED COST	30.13	584.61
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
CA CURCOTTER DATE	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.60	-33.20

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 12:40:52 ON 31 MAR 2003

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FILE 'REGISTRY' ENTERED AT 13:51:03 ON 01 APR 2003
               1 SEA ABB=ON "ETHYLENE OXIDE"/CN
 Ll
                 E PROPYLENE OXIDE/CN
               1 SEA ABB=ON "PROPYLENE OXIDE"/CN
 L2
                 E BUTYLENE OXIDE/RN
                 E BUTYLENE OXIDE/CN
L3
               1 SEA ABB=ON
                            "BUTYLENE OXIDE"/CN
L4
               O SEA ABB=ON L1 AND L2 AND L3
             110 SEA ABB=ON (75-21-8 AND 75-56-9 AND 26249-20-7)/CRN
                 E CHITIN/CN
               1 SEA ABB=ON CHITIN/CN
                 E AGAR/CN
L7
               1 SEA ABB=ON AGAR/CN
                E CARRAGEENAN/CN
              1 SEA ABB=ON CARRAGEENAN/CN
                 E ALGINATE/CN
                 E GUAR/CN
              1 SEA ABB=ON GUAR/CN
                E ARABIC/CN
                E GUM ARABIC/CN
              1 SEA ABB=ON "GUM ARABIC"/CN
L10
                E TRAGACANTH/CN
              1 SEA ABB=ON TRAGACANTH/CN
                E XANTHAN/CN
              1 SEA ABB=ON XANTHAN/CN
         635385 SEA ABB=ON L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR 12
ъ13.
              0 SEA ABB=ON (1398-61-4 AND 9002-18-0 AND 9000-07-1 AND
                9000-30-0 AND 9000-01-5 AND 9000-65-1 AND 11138-66-2)/CRN
L15
            680 SEA ABB=ON (1398-61-4 OR 9002-18-0 OR 9000-07-1 OR 9000-30-0
                OR 9000-01-5 OR 9000-65-1 OR 11138-66-2)/CRN
     FILE 'HCAPLUS' ENTERED AT 14:16:29 ON 01 APR 2003
L16
            498 SEA ABB=ON ?POLYSACCHARID?(3A)?ETHER? AND (?PROCES? OR
                ?TECHNIQ? OR ?PRODUC? OR ?PREP?)
L17
            112 SEA ABB=ON L16 AND (?ELECTRODIAL? OR ?SEMIPERMEAB? OR
                ?SEMI?(W)?PERMEAB? OR ?BIPOLAR? OR BI(W)?POLAR? OR L6 OR L7 OR
                L8 OR L9 OR L10 OR L11 OR L12 OR L15 OR ?CHITIN OR ?AGAR? OR
                ?CARRAGEEN? OR ?ALGINAT? OR ?GUAR? OR ?ARABIC? OR ?TRAGACANTH?
                OR ?XANTHAN?)
              6 SEA ABB=ON L17 AND (L1 OR L2 OR L3 OR L5) 6 cits in CA Rlus
L18
     FILE 'BIOSIS, CHEMREACT, CABA, BIOTECHDS, FSTA, JAPIO, JICST-EPLUS,
     PAPERCHEM2, WPIDS' ENTERED AT 14:28:06 ON 01 APR 2003
              O SEA ABB=ON L18 Ocition other databases
L19
     FILE 'HCAPLUS' ENTERED AT 14:32:52 ON 01 APR 2003
           9016 SEA ABB=ON ?CELLULOS?(3A)?ETHER? AND (?PROCES? OR ?TECHNIQ?
                OR ?PRODUC? OR ?PREP?)
             51 SEA ABB=ON L20 AND (?ELECTRODIAL? OR ?SEMIPERMEAB? OR
                ?SEMI?(W)?PERMEAB? OR ?BIPOLAR? OR BI(W)?POLAR?)
              7 SEA ABB=ON L21 AND (PH OR ?ELEC? (W) ?CURRENT?) Town on CA flux
     FILE 'BIOSIS, CHEMREACT, CABA, BIOTECHDS, FSTA, JAPIO, JICST-EPLUS,
    PAPERCHEM2, WPIDS' ENTERED AT 15:24:31 ON 01 APR 2003
2 SEA ABB=ON L22 2 Cits in other defabases.
L23
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=> d que stat 118
L1
              1 SEA FILE=REGISTRY ABB=ON
                                           "ETHYLENE OXIDE"/CN
L2
              1 SEA FILE=REGISTRY ABB=ON
                                           "PROPYLENE OXIDE"/CN
L3
              1 SEA FILE=REGISTRY ABB=ON
                                           "BUTYLENE OXIDE"/CN
            110 SEA FILE=REGISTRY ABB=ON
L5
                                           (75-21-8 AND 75-56-9 AND 26249-20-7)/
                CRN
1.6
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                                          CHITIN/CN
L7
              1 SEA FILE=REGISTRY ABB=ON AGAR/CN
1.8
              1 SEA FILE=REGISTRY ABB=ON CARRAGEENAN/CN
L9
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L11
              1 SEA FILE=REGISTRY ABB=ON
                                          TRAGACANTH/CN
L12
              1 SEA FILE=REGISTRY ABB=ON XANTHAN/CN
L15
            680 SEA FILE=REGISTRY ABB=ON
                                          (1398-61-4 OR 9002-18-0 OR 9000-07-1
                OR 9000-30-0 OR 9000-01-5 OR 9000-65-1 OR 11138-66-2)/CRN
L16
            498 SEA FILE=HCAPLUS ABB=ON ?POLYSACCHARID?(3A)?ETHER? AND
                (?PROCES? OR ?TECHNIQ? OR ?PRODUC? OR ?PREP?)
L17
            112 SEA FILE=HCAPLUS ABB=ON L16 AND (?ELECTRODIAL? OR ?SEMIPERMEAB
                ? OR ?SEMI?(W)?PERMEAB? OR ?BIPOLAR? OR BI(W)?POLAR? OR L6 OR
                L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR L15 OR ?CHITIN OR
                ?AGAR? OR ?CARRAGEEN? OR ?ALGINAT? OR ?GUAR? OR ?ARABIC? OR
                ?TRAGACANTH? OR ?XANTHAN?)
              6 SEA FILE=HCAPLUS ABB=ON L17 AND (L1 OR L2 OR L3 OR L5)
L18
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=> d ibib abs hitrn 1-6

L18 ANSWER 1 OF 6 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2003:43049 HCAPLUS

DOCUMENT NUMBER:

138:91690

TITLE:

Recycling of salts in the manufacture of modified

polysaccharide ethers from sodium

salt stock

INVENTOR(S):

Mallon, Charles B.; Vames, John S.; Sarlis, John I.;

See, Benito; Trampe, David M.; Datta, Rathin

PATENT ASSIGNEE(S): USA

SOURCE:

U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003013871	A1	20030116	US 2001-775760	20010202
PRIORITY APPLN. INFO.: AB Salts formed duri			2001-775760	20010202

Salts formed during the manuf. of polysaccharide ethers, e.g., sodium nitrate or sodium acetate, are converted to their corresponding acids and bases by means of an elec. current, preferably in combination with a bipolar membrane and suitable cation and/or anion membranes. The acids and bases recovered from the processes can be recycled, thereby avoiding the need to provide for disposal of the salts. Thus, a stream contg. .apprx.8% Na acetate, 0.5% NaOH, 4% glycols, and 0.5% cellulosic material recovered from a synthesis of hydroxyethyl cellulose was processed in a TS-2 ED stack contg. 6 cell pairs made up of AM-1, CM-2 and CURIUM ion-exchange membranes. The pH of the feed was raised to .apprx.11.5 by addn. of 40% NaOH whenever the pH dropped to .apprx.11. As a result of this pH control, .apprx.91% of the

```
acetate was transferred and cell resistance was kept under control.
      acetate concn. in the conc. was .apprx.20%. The product from
      the ED step was then processed through a column contg. Duolite
     C-467 to remove multivalent cations. Ca and Mg cations were below 1 ppm
      in the resulting stream. This product stream was then subjected
      to water splitting electrodialysis using a 2-compartment TS-2
     stack equipped with BP1, CM-1, and CMX ion exchange membranes. The stream
     processed with no difficulty and produced an acid/salt
     product contg. about 16% acetic acid and a sodium hydroxide
     product with a concn. of .apprx.10%. There was no evidence of
     irreversible membrane fouling and cell voltage remained low (.apprx.1.2
      V/cell pair) throughout the run.
     1398-61-4DP, Chitin, derivs. 9000-01-5DP, Gum
     Arabic, derivs. 9000-07-1DP, Carrageenan,
     derivs. 9000-30-0DP, Guar, derivs. 9000-65-1DP
      , Tragacanth, derivs. 9002-18-0DP, Agar,
     derivs. 11138-66-2DP, Xanthan gum, derivs.
     RL: IMF (Industrial manufacture); PREP (Preparation)
     (method for recycling of salts in manuf. of modified cellulose ether) 75-21-8, Ethylene oxide, reactions 75-56-9, Propylene
     oxide, reactions 26249-20-7, Butylene oxide
     RL: RGT (Reagent); RACT (Reactant or reagent)
         (method for recycling of salts in manuf. of modified cellulose ether)
L18 ANSWER 2 OF 6 HCAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                          2000:736368 HCAPLUS
DOCUMENT NUMBER:
                          134:163239
TITLE:
                          Synthesis of hydroxypropyl derivatives of
                          chitin and chitosan and observation of phase
                         behavior of their aqueous solutions
AUTHOR(S):
                         Asahina, Daisuke; Matsubara, Tomoyuki; Miyashita,
                         Yoshiharu; Nishino, Yoshiyuki
CORPORATE SOURCE:
                         Department of Material Systems Engineering, Tokyo
                         University of Agriculture and Technology, Koganei,
                         Tokyo, 184-8588, Japan
SOURCE:
                         Sen'i Gakkaishi (2000), 56(9), 435-442
                         CODEN: SENGA5; ISSN: 0037-9875
PUBLISHER:
                         Sen'i Gakkai
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Japanese
    Hydroxypropyl (HP) derivs. of chitin and chitosan with different
    degrees of substitution were synthesized via etherification of
    the polysaccharides with propylene oxide under various
    conditions. Mol. structure of the HP derivs. was characterized by 1H and
    13C NMR spectroscopy. On the basis of the spectral data, we proposed the
    equations to det. the molar substitution (MS), degree of pendants (DPs)
    and degree of substitution (DS) of the HP derivs. obtained. Phase
    behavior and liq.-cryst. characteristics of aq. solns. of the derivs. were
```

also examd. by light absorption and polarized optical microscopy (POM). Aq. solns. of highly substituted HP derivs. were phase-sepd. and became turbid at an elevated temp. The obsd. cloud point varied sensitively, depending on the degree of substitution and mol. wt. of HP derivs., concn. of the solns., and coexistence of inorg. salts. POM observations revealed that the formation of an optically anisotropic monophase prevailed in the concd. solns. of highly substituted HP derivs.

75-56-9DP, Propylene oxide, reaction products with chitin or chitosan 1398-61-4DP, Chitin, hydroxypropyl derivs.

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (synthesis of hydroxypropyl derivs. of chitin and chitosan

```
and observation of phase behavior of aq. solns.)
 ΤT
      75-56-9, Propylene oxide, reactions 1398-61-4,
       Chitin
      RL: RCT (Reactant); RACT (Reactant or reagent)
          (synthesis of hydroxypropyl derivs. of chitin and chitosan
          and observation of phase behavior of aq. solns.)
 L18 ANSWER 3 OF 6 HCAPLUS COPYRIGHT 2003 ACS
 ACCESSION NUMBER:
                            1999:708699 HCAPLUS
 DOCUMENT NUMBER:
                            131:326313
 TITLE:
                            Etherified polysaccharides as
                            concrete modifiers for improved workability
 INVENTOR (S):
                            Veen, Uko; Lamberti, Vincent Joseph Marie Alphonse;
                            Bleeker, Ido Pieter
 PATENT ASSIGNEE(S):
                            Cooperatieve Verkoop- en Productievereniging Van
                            Aardappelmeel en Derivaten, Neth.
 SOURCE:
                            PCT Int. Appl., 47 pp.
                            CODEN: PIXXD2
 DOCUMENT TYPE:
                            Patent
 LANGUAGE:
                            English
 FAMILY ACC. NUM. COUNT:
 PATENT INFORMATION:
      PATENT NO.
                         KIND DATE
                                                APPLICATION NO. DATE
                                                -----
      WO 9955632
                                               WO 1999-NL248
                         A1
                               19991104
                                                                  19990427
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              DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
              MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
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              CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
      EP 955277
                         A1 19991110
                                              EP 1998-201379
                                                                19980428
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      AU 9935397
                              19991116
                         A1
                                               AU 1999-35397
                                                                  19990427
      EP 1080048
                         A1
                              20010307
                                               EP 1999-917238
                                                                  19990427
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO
PRIORITY APPLN. INFO.:
                                            EP 1998-201379
                                                            A 19980428
                                                              W 19990427
                                            WO 1999-NL248
AB
     Polysurfactant-like polysaccharide concrete modifier preferably derived
     from potato starch is used to improve workability of concrete mixes. The
     manuf. of the modifier is described in details and includes
     etherification of polysaccharides up to a degree of
     substitution allowing dissolving the modifier in a 95 wt.% ethanol soln.
IΤ
     75-21-8, Oxirane, processes 75-56-9,
     processes 9000-30-0, Guar 26249-20-7
     , Butylene oxide
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
         (etherified polysaccharides as concrete modifiers
        for improved workability)
REFERENCE COUNT:
                                  THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L18 ANSWER 4 OF 6 HCAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                           1992:429148 HCAPLUS
```

DOCUMENT NUMBER:

117:29148

TITLE:

Hydrophobically modified hydroxybutyl ethers of

polygalactomannan

INVENTOR(S):

Zody, George M.; Morgan, Michael E.

PATENT ASSIGNEE(S):

Hi-Tek Polymers, Inc., USA Can. Pat. Appl., 21 pp.

SOURCE:

CODEN: CPXXEB

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CA 2023215	AA	19911230	CA 1990-2023215	19900814
CA 2023215	C	20010327		
US '5233032	A	19930803	US 1990-546560	19900629
RIORITY APPLN. INFO.:		US	1990-546560 A	19900629
P The title others	usofu	l as thickeners	and can be used	21000 00

PR AB The title ethers, useful as thickeners and can be used alone or with other surfactants in drilling muds, are prepd. to contain hydrophobic groups such as C8-28 alkoxy, HOZO (Z = C8-28 alkylene bearing OH group on the C beta to ether O), and R1OCH2CH(OH)CH2O (R1 = C5-25 alkyl) wherein the mol. hydrophilic substitution degree is 0.2-1, and the mol. hydrophobic substitution degree is 0.001-0.2. Thus, a hydroxybutylated guar was prepd. in a customary way using butylene oxide,. combined (90 parts) with iso-PrOH 130, ammonium lauryl sulfate (I) 2, water 36, and 1,2-epoxyhexadecane 18, stirred, heated to 70.degree. and combined with KOH 3.15 parts, and heated at 70.degree. for addnl. 3 h to give a product, 400 parts of a 0.5% aq. soln. of which at pH 6.0 with a 28% aq. soln. of I in 0.2, 0.4, 06, and 0.8 part showed Brookfield viscosity of 905, 4150, 6680, and 6900, resp.

26249-20-7, Butylene oxide

RL: USES (Uses)

(hydroxyalkylation with, of galactomannan) 102962-18-5DP, etherified to hydrophobic groups

RL: PREP (Preparation)

(prepn. of amphiphilic, for thickeners)

L18 ANSWER 5 OF 6 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1992:410198 HCAPLUS

DOCUMENT NUMBER:

117:10198

TITLE:

ΙT

Process for preparing

hydrophobically modified guar ethers Zody, George M.; Morgan, Michael E.

PATENT ASSIGNEE(S):

Hi-Tek Polymers, Inc., USA

SOURCE:

Can. Pat. Appl., 19 pp.

CODEN: CPXXEB

DOCUMENT TYPE:

Patent

LANGUAGE:

INVENTOR(S):

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CA 2023324	AA	19920106	CA 1990-2023324	19900815
PRIORITY APPLN. INFO.			JS 1990-547968	
			s are prepd . by a	
derivatization r	rocess	in which the	1st step is carrie	d out with
hydrophilic read	ents (e.g. alkylene	oxide) under alk.	conditions, and,

without isolation of the products, the 2nd step carried out with

hydrophobic reagents (e.g. oxirane compds.) dissolved in org. solvents and reacted with the guar still in particulate form. In this manner, NaOH-pre-activated guar splits were hydroxyalkylated with propylene oxide, and derivatized with 1,2-epoxyhexadecane in propylene oxide to give an amphiphilic ether which exhibited high viscosity in aq. solns. contg. a surfactant.
75-56-9DP, Propylene oxide, galactomannan mixed etherified with

9000-30-0DP, Guar gum, etherified with alkylene oxides and epoxy compds. 26249-20-7DP, Butylene oxide, galactomannan mixed etherified with

RL: PREP (Preparation)

(prepn. of amphiphilic, for thickeners)

L18 ANSWER 6 OF 6 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1992:8169 HCAPLUS

DOCUMENT NUMBER:

116:8169

TITLE:

ΙT

Process for the preparation of

hydroxyalkyl ethers of

polysaccharides

INVENTOR(S):

Srivastava, Harish Chandra; Phadnis, Shashikant

Purushottam; Parikh, Bharat Siddharthbhai

PATENT ASSIGNEE(S):

Ahmedabad Textile Industry's Research Assoc., India

SOURCE:

Indian, 12 pp.

CODEN: INXXAP

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE PATENT NO. APPLICATION NO. DATE ---------IN 166309 Α 19900407 IN 1987-BO207 19870702 PRIORITY APPLN. INFO.: IN 1987-B0207 19870702 The title process, without use of expensive org. solvents, comprises hydroxyalkylation of polysaccharides, e.g. starch and gums in aq. or aq.-org. mixed media at 40-60.degree., in the presence of alk. catalyst and a gelling suppressant, e.g. inorg. salts. Thus, adding starch 100 to a soln. of Na2SO4 45 in water 150, flushing with N, adding

Ca(OH)2 5, and then ethylene oxide 8 parts, and heating to 45 .+-. 2.degree. for 3 h gave a starch ether with degree of substitution 0.28. The polysaccharide derivs. are useful as sizes and adhesives for textiles, fibers and paper.

TT 39465-11-7P, Hydroxyethyl guar gum

RL: PREP (Preparation)

(prepn. of, gelling prevention in)

IT 75-21-8, Ethylene oxide, uses

RL: USES (Uses)

(reagents, for hydroxyalkylation of polysaccharides)



01/04/2003

=> d que stat 122

9016 SEA FILE=HCAPLUS ABB=ON ?CELLULOS?(3A)?ETHER? AND (?PROCES? OR ?TECHNIQ? OR ?PRODUC? OR ?PREP?) L20

51 SEA FILE=HCAPLUS ABB=ON L20 AND (?ELECTRODIAL? OR ?SEMIPERMEAB L21

? OR ?SEMI?(W)?PERMEAB? OR ?BIPOLAR? OR BI(W)?POLAR?)

7 SEA FILE=HCAPLUS ABB=ON L21 AND (PH OR ?ELEC?(W)?CURRENT?) 1.22

=> d 122 ibib abs hitrn 1-7

L22 ANSWER 1 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

2003:43049 HCAPLUS

DOCUMENT NUMBER:

138:91690

TITLE:

Recycling of salts in the manufacture of modified

polysaccharide ethers from sodium salt stock

INVENTOR (S):

Mallon, Charles B.; Vames, John S.; Sarlis, John I.;

See, Benito; Trampe, David M.; Datta, Rathin

PATENT ASSIGNEE(S):

SOURCE:

U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ---------US 2003013871 A1 US 2001-775760 20030116 US 2001-775760 20010202 US 2001-775760 20010202

PRIORITY APPLN. INFO.: . Salts formed during the manuf. of polysaccharide ethers, e.g., sodium nitrate or sodium acetate, are converted to their corresponding acids and bases by means of an elec. current, preferably in combination with a bipolar membrane and suitable cation and/or anion membranes. The acids and bases recovered from the processes can be recycled, thereby avoiding the need to provide for disposal of the salts. Thus, a stream contg. .apprx.8% Na acetate, 0.5% NaOH, 4% glycols, and 0.5% cellulosic material recovered from a synthesis of hydroxyethyl cellulose was processed in a TS-2 ED stack contg. 6 cell pairs made up of AM-1, CM-2 and CURIUM ion-exchange membranes. The pH of the feed was raised to .apprx.11.5 by addn. of 40% NaOH whenever the pH dropped to .apprx.11. As a result of this pH control, .apprx.91% of the acetate was transferred and cell resistance was kept under control. Na acetate concn. in the conc. was .apprx.20%. The product from the ED step was then processed through a column contg. Duolite C-467 to remove multivalent cations. Ca and Mg cations were below 1 ppm in the resulting stream. This product stream was then subjected to water splitting electrodialysis using a 2-compartment TS-2 stack equipped with BP1, CM-1, and CMX ion exchange membranes. The stream processed with no difficulty and produced an acid/salt product contg. about 16% acetic acid and a sodium hydroxide product with a concn. of .apprx.10%. There was no evidence of irreversible membrane fouling and cell voltage remained low (.apprx.1.2 V/cell pair) throughout the run.

L22 ANSWER 2 OF 7 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2000:117084 HCAPLUS

DOCUMENT NUMBER:

132:153513

TITLE:

Production of polysaccharide hydroxyalkyl

ethers

```
INVENTOR(S):
                           Mallon, Charles B.; Vames, John S.; Sarlis, John
                           Ioannis; See, Benito; Trampe, David M.; Datta, Rathin
 PATENT ASSIGNEE(S):
                           Union Carbide Chemicals & Plastics Technology Corp.,
                           USA
 SOURCE:
                           PCT Int. Appl., 32 pp.
                           CODEN: PIXXD2
 DOCUMENT TYPE:
                           Patent
 LANGUAGE:
                           English
 FAMILY ACC. NUM. COUNT:
 PATENT INFORMATION:
      PATENT NO.
                       KIND DATE
                                             APPLICATION NO. DATE
      -----
                              -----
      WO 2000008059
                       A1 20000217
                                             WO 1999-US17597 19990803
          W: AL, AU, BA, BB, BG, BR, CA, CN, CU, C2, EE, GE, HU, ID, IL, IS, JP, KR, KZ, LC, LK, LR, LT, LV, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY,
              KG, KZ, MD, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
              ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
              CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
      CA 2339374
                       AA
                              20000217
                                            CA 1999-2339374 19990803
                              20000228
      AU 9954643
                        A1
                                            AU 1999-54643
                                                               19990803
      EP 1109834
                                        EP 1999-940869 19990803
                              20010627
                        A1
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
                              20011009
                   A
                                             BR 1999-12637
                                                               19990803
 PRIORITY APPLN. INFO.:
                                          US 1998-95263P P 19980804
                                          WO 1999-US17597 W 19990803
AB
      Processes for the prodn. of polysaccharide
      ethers, e.g. cellulose hydroxyethyl ether, are
      disclosed wherein salts formed after the swelling and neutralization of
      the process, e.g., sodium nitrate or sodium acetate, are
      converted to their corresponding acids and bases by means of an
      elec. current, preferably in combination with a
     bipolar membrane and suitable cation and/or anion membranes.
     acids and bases recovered from the processes can be recycled,
      thereby avoiding the need to provide for disposal of the salts.
REFERENCE COUNT:
                          8
                                THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L22 ANSWER 3 OF 7 HCAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                          1985:191189 HCAPLUS
DOCUMENT NUMBER:
                          102:191189
TITLE:
                          Oral device for osmotic delivery of drugs
INVENTOR(S):
                          Edgren, David; Wong, Patrick S. L.; Theeuwes, Felix
PATENT ASSIGNEE(S):
                          Alza Corp., USA
SOURCE:
                          U.S., 9 pp.
                          CODEN: USXXAM
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
                   KIND DATE
     PATENT NO.
                                            APPLICATION NO. DATE
                      ----
                             -----
     US 4503030
                       Α
                             19850305
                                            US 1983-501573
                                                              19830606
     US 4587117
                      Α
                             19860506
                                            US 1985-693649
                                                              19850122
PRIORITY APPLN. INFO.:
                                         US 1983-501573
                                                              19830606
    An oral osmotic delivery device for dispensing a drug to certain
```

pH regions of the gastrointestinal tract consists of a wall formed of a semipermeable pH-sensitive compn. that surrounds a compartment contg. a drug, with a passage through the wall connecting the exterior of the device with the compartment. The device delivers the drug at a controlled rate in the region of the gastrointestinal tract having a pH <3.5 (stomach), self-destructs, and releases the remainder of the drug in the region of the gastrointestinal tract having a pH >3.5 (small intestine) thereby providing total availability for drug absorption. The wall is formed of a semipermeable material, such as a cellulose ester or ether, blended with a pH-sensitive material, such as a cellulose carboxylic acid ester which keeps its integrity at pH 1.0-3.58 but loses it at pH >3.5, and optionally a flux enhancer, such as hydroxypropyl Me cellulose [9004-65-3]. Thus, a device for delivering hydralazine-HCl [304-20-1] was manufd. by prepg. cores (275 mg) contg. drug 18.2, mannitol 75.9, hydroxypropyl Me cellulose 2.9, and stearic acid 3% by wt. and then coating the core with a pH -sensitive material consisting of cellulose acetate [9004-35-7] and hydroxypropyl Me cellulose phthalate [9050-31-1] coated to a wall wt. .apprx.20 mg and air-dried at 50.degree. for 48 h, after which a 10 mill hole was laser-drilled through the wall.

L22 ANSWER 4 OF 7 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1985:12091 HCAPLUS

DOCUMENT NUMBER:

102:12091

TITLE:

Elimination of acid-base generation ('water-splitting') in electrodialysis

AUTHOR (S):

Rubinstein, I.; Warshawsky, A.; Schechtman, L.; Kedem,

CORPORATE SOURCE:

SOURCE:

Weizmann Inst. Sci., Rehovot, Israel Desalination (1984), 51(1), 55-60 CODEN: DSLNAH; ISSN: 0011-9164

DOCUMENT TYPE:

LANGUAGE:

Journal English

Membranes contg. a crown ether were prepd. The membranes become pos. charged by complexing alkali-metal ions. In these anion-exchange membranes, not contg. amino groups, pH changes caused by above-limiting currents were very small in contrast to the substantial acid-base generation by conventional A membranes. This is consistent with R. Simons's model (1979). It is suggested that both suppression of acidification and the dynamic nature of the pos. charges may help to avoid fouling.

L22 ANSWER 5 OF 7 HCAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER:

DOCUMENT NUMBER:

1976:136377 HCAPLUS

TITLE:

84:136377

Preparation and properties of polymer

sorbents containing p-aminomethylphenylboric acid

groups

AUTHOR(S):

Kolodkina, I. I.; Val'kovskii, D. G.; Pichuzhkina, E. I.; Ivanova, E. A.; Rogozhin, S. V.; Yurkevich, A. M.

CORPORATE SOURCE: SOURCE:

Vses. Nauchno-Issled. Vitam. Inst., Moscow, USSR Vysokomolekulyarnye Soedineniya, Seriya A (1976),

18(1), 47-52 CODEN: VYSAAF; ISSN: 0507-5475

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

The title sorbents were obtained by reaction of chloromethylated and aminated divinylbenzene-styrene copolymer (I) [9003-70-7] with [p-(bromomethyl)phenyl]boroxole (II) [51239-44-2] yielding polymer contg. groups R1CH2N+R22CH2C6H4B-(OH)3 (R1 = polymer residue, R2 = Me or Et), or by analogous reactions of (diethylamino)ethyl ethers of Sephadexes and cellulose or polyethylenepolyamine with II. Sorption properties of the products derived from I, Sephadex A-25 (diethylamino)ethyl ether [57866-54-3], and Sephadex A-50 (diethylamino)ethyl ether [39455-31-7] with resp. to adenosine 5'-phosphomorpholide [7331-13-7] or adenosine phosphates indicated that the bipolar structure of the sorbents facilitated formation of complexes at a wide range of pH.

L22 ANSWER 6 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER:

1966:37056 HCAPLUS

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: 64:37056 64:6923d-h

TITLE:

Interaction of synthetic high-molecular-weight

materials with sulfate and organic salts of aluminum

Kirakos'yants, M. Kh.; Strakhov, I. P.

AUTHOR(S): SOURCE:

Nauchn. Tr. Mosk. Tekhnol. Inst. Legkoi Prom. (1964),

No. 30, 26-33

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

The interaction of various synthetic polymeric materials, having different functional groups, with sulfate-type Al salts was investigated by the method of equil. dialysis (Chanutin, et al., CA 36, 45315). The polymers tested were: poly(vinyl alc.) (PVA), mol. wt. 60,000; poly(acrylic acid) (PAA), mol. wt. 80,000; copolymer of styrene and maleic anhydride (S-MA), mol. wt. not detd.; urea-HCHO resin (U-F), mol. wt. not detd.; and Na carboxymethyl cellulose (CMC), mol. wt. 243,000, degree of etherification These polymers have functional groups similar to those in collagen, and can form coordination bonds with Al. In tanning, these polymers enhance fixation of Al complexes in hides. Aq. solns. (20 g./l.) of polymers were used. At first, an aq. soln. of Al2(SO4)3.18H2O (brought to pH 4.2) was used; various salts of org. acids were then added. In all these solns., Al concn. corresponded to 20 g./l. The amts. of org. salts were 1.0-1.5 g.-equiv. per 1.0 g.-atom Al. The pH changed (4.0-4.7). The salts were tartrate, citrate, oxalate, and lactate. Solns. of polymers and Al salts were placed in a dialyzer with a cellophane film as a semipermeable membrane. As Al complex diffused into the cell with polymer soln., the latter turned gradually into a gel. Strongly coordinated bonds were formed with functional groups of polymers. The remaining (unbound) Al attained equil. on both sides of the membrane. The amt. of bound Al was detd. from the initial and later concns. of Al salt. The amt. of bound Al was higher when org. salts were present than in the case of nonstabilized Al sulfate. The enhancement of Al binding followed the order: S-MA > PVA > U-F > PAA > CMC, and for salt-stabilized sulfates: PVA tartrate > citrate > lactate > oxalate; PAA tartrate > citrate > oxalate > lactate; U-F citrate > tartrate > lactate > oxalate; CMC citrate > tartrate > oxalate > lactate; and S-MA tartrate > citrate > lactate > oxalate. The polymeric gels formed apparently had a cross-linked structure; the retained Al could not be removed by water. contrast, nonstabilized Al sulfate did not interact strongly with functional groups of polymers, since Al compds. were leached out with water.

L22 ANSWER 7 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

1934:6210 HCAPLUS

ORIGINAL REFERENCE NO.:

28:6210

TITLE:

28:799d-f Method of electrodialysis. Serum

electrodialysis with glycine membranes

AUTHOR(S): SOURCE:

Ettisch, G.; de Loureiro, J. A. Biochem. Z. (1933), 266, 422-35

DOCUMENT TYPE: LANGUAGE:

Journal Unavailable

AB Membranes are prepd. from collodion contg. glycine, 1:20,000, which decreases the viscosity. Add 2 mg. powd. glycine to 50 cc. collodion, shake for a few min. and pour on a glass plate after standing about 1/2 hr. The glycine suspension should be preferably fresh. Electrodialysis proceeds very rapidly with these membranes because they are permeable to anions, and after 35 min. the pH reaches a definitive value. Because of the shortness of time during which the pH drops from 7.8 to 6.8, usually about 10 min., a large part of the globulin is pptd. quickly. If a dry glycine-collodion membrane is used at the anode the end pH value is not attained so rapidly. It does not affect the results whether a parchment or cellulose membrane is used at the cathode.

```
=> d que stat 123
 L20
            9016 SEA FILE=HCAPLUS ABB=ON ?CELLULOS?(3A)?ETHER? AND (?PROCES?
                 OR ?TECHNIQ? OR ?PRODUC? OR ?PREP?)
              51 SEA FILE=HCAPLUS ABB=ON L20 AND (?ELECTRODIAL? OR ?SEMIPERMEAB
 L21
                 ? OR ?SEMI?(W)?PERMEAB? OR ?BIPOLAR? OR BI(W)?POLAR?)
               7 SEA FILE=HCAPLUS ABB=ON L21 AND (PH OR ?ELEC?(W)?CURRENT?)
 L22
 L23
               2 SEA L22
 => d ibib abs 123 1-2
 L23 ANSWER 1 OF 2 WPIDS (C) 2003 THOMSON DERWENT
 ACCESSION NUMBER:
                       2001-112499 [12]
 CROSS REFERENCE:
                       2001-091751 [10]
 DOC. NO. CPI:
                       C2001-033517
 TITLE:
                       Method for controlling the flux of penetrants across an
                       adaptable semi-permeable barrier is
                       useful for administering an agent to a mammalian body or
                       a plant and for generating an immune response by
                       vaccinating the mammal.
 DERWENT CLASS:
                      A18 A28 A96 B05 B07 D16 D22
 INVENTOR(S):
                      CEVC, G; RICHARDSEN, H; WEILAND-WAIBEL, A;
                      WEILAND-WEIBEL, A
PATENT ASSIGNEE(S):
                       (IDEA-N) IDEA AG
COUNTRY COUNT:
                       95
PATENT INFORMATION:
     PATENT NO KIND DATE
                               WEEK
                                         LA
                                              PG
     WO 2001001963 A1 20010111 (200112)* EN 110
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM
            DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC
            LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE
            SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
     AU 2000061557 A 20010122 (200125)
     BR 2000012178 A 20020312 (200226)
     EP 1189598
                 A1 20020327 (200229) EN
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI
     CZ 2002000038 A3 20020515 (200241)
     CN 1359288
                A 20020717 (200268)
     HU 2002001454 A2 20021228 (200308)
    JP 2003503442 W 20030128 (200309)
                                             109
APPLICATION DETAILS:
```

PATENT NO KI	ND	APPLICATION	DATE
WO 2001001963 AU 2000061557 BR 2000012178	A A	WO 2000-EP6367 AU 2000-61557 BR 2000-12178	20000705 20000705 20000705
EP 1189598	A1	EP 2000-947939	20000705 20000705
CZ 2002000038	A3	WO 2000-EP6367	20000705 20000705
CN 1359288 HU 2002001454	A No	CN 2000-809916	20000705 20000705 20000705

JP 2003503442 W

HU 2002-1454 20000705 WO 2000-EP6367 20000705 JP 2001-507458 20000705

FILING DETAILS:

PATENT NO KIND	PATENT NO
AU 2000061557 A Based on	WO 200101963
BR 2000012178 A Based on	WO 200101963
EP 1189598 Al Based on	WO 200101963
CZ 2002000038 A3 Based on	WO 200101963
HU 2002001454 A2 Based on	WO 200101963
JP 2003503442 W Based on	WO 200101963

PRIORITY APPLN. INFO: WO 1999-EP4659 19990705

2001-112499 [12] WPIDS

2001-091751 [10] CR

AB

WO 200101963 A UPAB: 20030206

NOVELTY - A method for controlling the flux of penetrants across an adaptable semi-permeable porous barrier is new.

DETAILED DESCRIPTION - A method for controlling the flux of penetrants across an adaptable semi-permeable membrane comprises suspending the penetrants in a polar liquid in the form of fluid droplets surrounds by a membrane-like coating comprising at least two kinds of amphiphilic substances with a tendency to aggregate, selecting a dose of the penetrants to control the flux of the penetrants across the barrier and applying the selected dose of the formulation onto the area of the barrier. The amphiphilic substances differ by a factor of at least 10 in solubility in the polar liquid and the homo-aggregates of the more soluble substance and hetero-aggregates have a preferred average diameter smaller than the diameter of the homo-aggregates of the less soluble substance. The more soluble substance tends to solubilize the droplet and comprises up to 99% of the solubilizing concentration or saturating concentration in the unstabilized droplet. The presence of the more soluble substance lowers the average elastic energy of the coating by at least 5 times preferably more than 10 times the average elastic energy of red blood cells or of phospholipid bilayers with fluid aliphatic chains. The penetrants are able to transport agents through the pores of the barrier or enable agent permeation through the pores after the penetrants have entered the pores.

INDEPENDENT CLAIMS are included for:

- (i) a kit containing the formulation;
- (ii) a patch containing the formulation; and
- (iii) a method of administering an agent to a mammalian body or plant comprising the novel method.

 USE - The method is useful for administering an agent to a mammalian

body or a plant, for generating an immune response by vaccinating the mammal and for treating inflammatory disease, dermatosis, kidney or liver failure, adrenal insufficiency, aspiration syndrome, Behcet syndrome, bites and stings, blood disorders (cold-hemagglutinin disease), hemolytic anaemia, hypereosinophilic, hypoplastic anaemia, macroglobulinaemia and thrombocytopenic purpura), bone disorders, cerebral oedema, Cogan's syndrome, congenital adrenal hyperplasia, connective tissue disorders (lichen, lupus erythematosus, polymyalgia rheumatica, polymyositis and dermatomyositis), epilepsy, eye disorders (cataracts), Graves' ophthalmopathy, hemangioma, herpes infections, neuropathies, retinal vasculitis, scleritis, gastro-intestinal disorders (inflammatory bowel disease, nausea and oesophageal damage), hypercalcaemia, infections, Kawasaki disease, myasthenia gravis, pain syndromes, polyneuropathies,

pancreatitis, respiratory disorders (asthma), rheumatoid disease, osteoarthritis, rhinitis, sarcoidosis, skin diseases, alopecia, eczema, erythema multiforme, lichen, pemphigus and pemphigoid, psoriasis, pyoderma gangrenosum, urticaria and thyroid and vascular disorders.

ADVANTAGE - Increasing the applied dose above a threshold level affects both the drug/penetrant distribution and also determines the rate of penetrant transport across the barrier. Dwg.0/14

L23 ANSWER 2 OF 2 WPIDS (C) 2003 THOMSON DERWENT

ACCESSION NUMBER:

2000-205675 [18] WPIDS

DOC. NO. CPI:

C2000-063455

TITLE:

Production of polysaccharide ether, for

industrial and personal care applications, includes

subjecting salts to electric current.

DERWENT CLASS:

INVENTOR(S):

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COUNTRY COUNT: 75

PATENT INFORMATION:

PATENT N	O KIND	DATE	WEEK	LA	PG

WO 2000008059 A1 20000217 (200018)* EN 32

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW

W: AL AU BA BB BG BR CA CN CU CZ EE GE HU ID IL IS JP KR KZ LC LK LR LT LV MG MK MN MX NO NZ PL RO RU SG SI SK SL TR TT UA US UZ VN YU 7.A

AU 9954643 A 20000228 (200030)

EP 1109834 A1 20010627 (200137) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

BR 9912637

A 20011009 (200168) A 20011114 (200217) CN 1322215

MX 2001001247 A1 20010501 (200227)

APPLICATION DETAILS:

PA	TENT NO K	IND	AP	PLICATION	DATE
ΑU	2000008059 9954643 1109834	A1 A A1	ΑU	1999-US17597 1999-54643 1999-940869	19990803 19990803 19990803
BR	9912637	A	WO BR	1999-US17597 1999-12637 1999-US17597	19990803 19990803
	1322215 2001001247	A A1	CN	1999-811749 2001-1247	19990803 19990803 20010201

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 9954643	A Based on	WO 200008059
EP 1109834	Al Based on	WO 200008059
BR 9912637	A Based on	WO 200008059

PRIORITY APPLN. INFO: US 1998-95263P 19980804 AN 2000-205675 [18] WPIDS AB WO 200008059 A UPAR 2000

WO 200008059 A UPAB: 20000412

NOVELTY - Polysaccharide ether, is produced by subjecting the salts to an electric current effective to promote the conversion of the salts to their respective acids and bases.

DETAILED DESCRIPTION - Production of polysaccharide ether comprises:

(i) treating a polysaccharide with a basic compound to promote swelling of the polysaccharide,

(ii) reacting the polysaccharide with at least one derivatizing agent in a liquid medium to promote reaction between the polysaccharide and the derivatizing agent and form a reaction product comprising a polysaccharide ether and the basic compound,

(iii) treating at least a portion of the reaction product with an acidic compound, to provide a neutralizing liquid comprising a salt of the acidic compound and the basic compound, and

(iv) separating the polysaccharide ether from at least one of the reaction product of the neutralized liquid.

The improvement comprises subjecting the neutralized liquid to an electric current and suitable means to promote the conversion of the salt to the acidic compound and the basic compound.

USE - For the production of polysaccharide ethers, especially hydroxyethyl cellulose, for use in industrial application, e.g., viscosity adjuster, suspension aids and oil field drilling, and personal care application, e.g., ointments, skin creams, lotions and soaps.

ADVANTAGE - The acids and bases recovered from the processes can be recycled, thus avoiding the need to provide for disposal of the salts. The degree of fouling of the membrane can be reduced and the ionic mobility of the salt through the membrane can be enhanced by conducting the electrodialysis at an alkaline pH.

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